The mission of the Spina Bifida Association of Greater New England is to build a better and brighter future for all those impacted by Spina Bifida.

SBAGNE is supported by generous individual donors, matching gifts, grants, corporate sponsors, and our fundraising events.

You can help make a difference in the Spina Bifida community today by making a personal donation, corporate contribution, or by getting involved in one of our various fundraising programs.

To learn more about ways you can give, please visit our website.

Phone: 888-479-1900
Email: Info@SBAGreaterNE.org
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Affiliated with the Spina Bifida Association (SBA)
For more information go to spinabifidaassociation.org

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SUPPORT AND EDUCATION FOR THOSE IMPACTED BY SPINA BIFIDA
ABOUT SPINA BIFIDA

Spina Bifida is a neural tube defect that happens in the first month of pregnancy when the spinal column doesn’t close completely.

Those who are born with Spina Bifida experience conditions such as paralysis, bladder and bowel difficulties, learning disabilities, depression, and the risk of latex allergy.

Spina Bifida has no known cause and each of the 65 million women of childbearing age in this country is at risk for a Spina Bifida affected pregnancy.

Since 1971, the Spina Bifida Association of Greater New England has served those living with Spina Bifida, their families and supporters.

SBAGNE has accomplished much, but much remains to be done. We are continually working to identify new and resourceful ways to support and enrich the lives of those living with Spina Bifida.

WHAT WE DO

- Resources, information and referral
- Educational programs
- Teen transition programs
- Parent to Parent Program
- Financial assistance
- Newsletter and social media updates
- Spina Bifida advocacy
- Social and community building events
- Monthly adult group meetings
- Youth and family programs